

S-S Synapsis during Class Switch Recombination Is Promoted by Distantly Located Transcriptional Elements and Activation-Induced Deaminase

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SUMMARY

Molecular mechanisms underlying synapsis of activation-induced deaminase (AID)-targeted S regions during class switch recombination (CSR) are poorly understood. By using chromosome conformation capture techniques, we found that in B cells, the E μ and 3'E α enhancers were in close spatial proximity, forming a unique chromosomal loop configuration. B cell activation led to recruitment of the germline transcript (GLT) promoters to the E μ :3'E α complex in a cytokine-dependent fashion. This structure facilitated S-S synapsis because S μ was proximal to E μ and a downstream S region was corecruited with the targeted GLT promoter to E μ :3'E α . We propose that GLT promoter association with the E μ :3'E α complex creates an architectural scaffolding that promotes S-S synapsis during CSR and that these interactions are stabilized by AID. Thus, the S-S synaptosome is formed as a result of the self-organizing transcription system that regulates GLT expression and may serve to guard against spurious chromosomal translocations.

INTRODUCTION

In mature B cells, class switch recombination (CSR) promotes diversification of effector functions encoded in constant (C μ) regions while maintaining the original antigen binding specificity arising from V(D)J joining. The mouse IgH locus (*Igh*) contains eight C μ genes (encoding μ , δ , γ 3, γ 1, γ 2a, γ 2b, ϵ , and α chains) that are embedded in a 220 kb tract (Chevallard et al., 2002; Zhou et al., 2002) and are located downstream of the V, D, and J segments. Isotype class switching involves an intrachromosomal

deletional rearrangement that focuses on regions of 1–10 kb of repetitive switch (S) DNA located upstream of each C μ gene (with the exception of C δ). In class switch recombination (CSR), hybrid S μ -S α junctions are formed on the chromosome while the intervening genomic material is looped out and excised as a circle. CSR is targeted to specific S regions by the selective transcriptional activation of germline transcripts (GLTs) (reviewed in Manis et al., 2002; Stavnezer, 2000) and may be dependent on isotype-specific factors (Kenter et al., 2004; Ma et al., 2002; Shanmugam et al., 2000). Activation-induced deaminase (AID) is a single-strand DNA-specific deaminase that is required for CSR (Muramatsu et al., 2000) and leads to the formation of double-strand breaks (DSBs) in S regions (Catalan et al., 2003; Rush et al., 2004; Schrader et al., 2005; Wuerffel et al., 1997). AID- and uracil-DNA glycosylase (UNG)-dependent DSBs are processed through a cascade of events mediated by base excision repair (BER), mismatch repair, nonhomologous end joining (NHEJ), and error-prone polymerases, thereby facilitating the formation of S-S junctions (Dudley et al., 2005; Kenter, 2005).

S regions targeted for recombination can be separated by as much as 150 kb, and this distance is likely to be an impediment for S-S synapse formation. It has been inferred that two DNA repair proteins, 53BP1 and γ -H2AX, are involved in S region synapsis because targeted deletion of the genes encoding these proteins abolished or reduced CSR, respectively, whereas intra-S μ rearrangements and mutations were retained (Reina-San-Martin et al., 2003, 2007). However, 53BP1 and γ -H2AX normally engage in DSB repair processes that protect against large chromosomal deletions (Bekker-Jensen et al., 2005; Celeste et al., 2002; Mochan et al., 2004; Rogakou et al., 1999). This leads to the perplexing question of how distantly located S region-specific DSBs are recruited to partner in a CSR reaction that leads to intrachromosomal deletion.

The mouse β -globin genes and their locus control region (LCR) located more than 50 kb away engage in higher-order loop structures during transcription, and

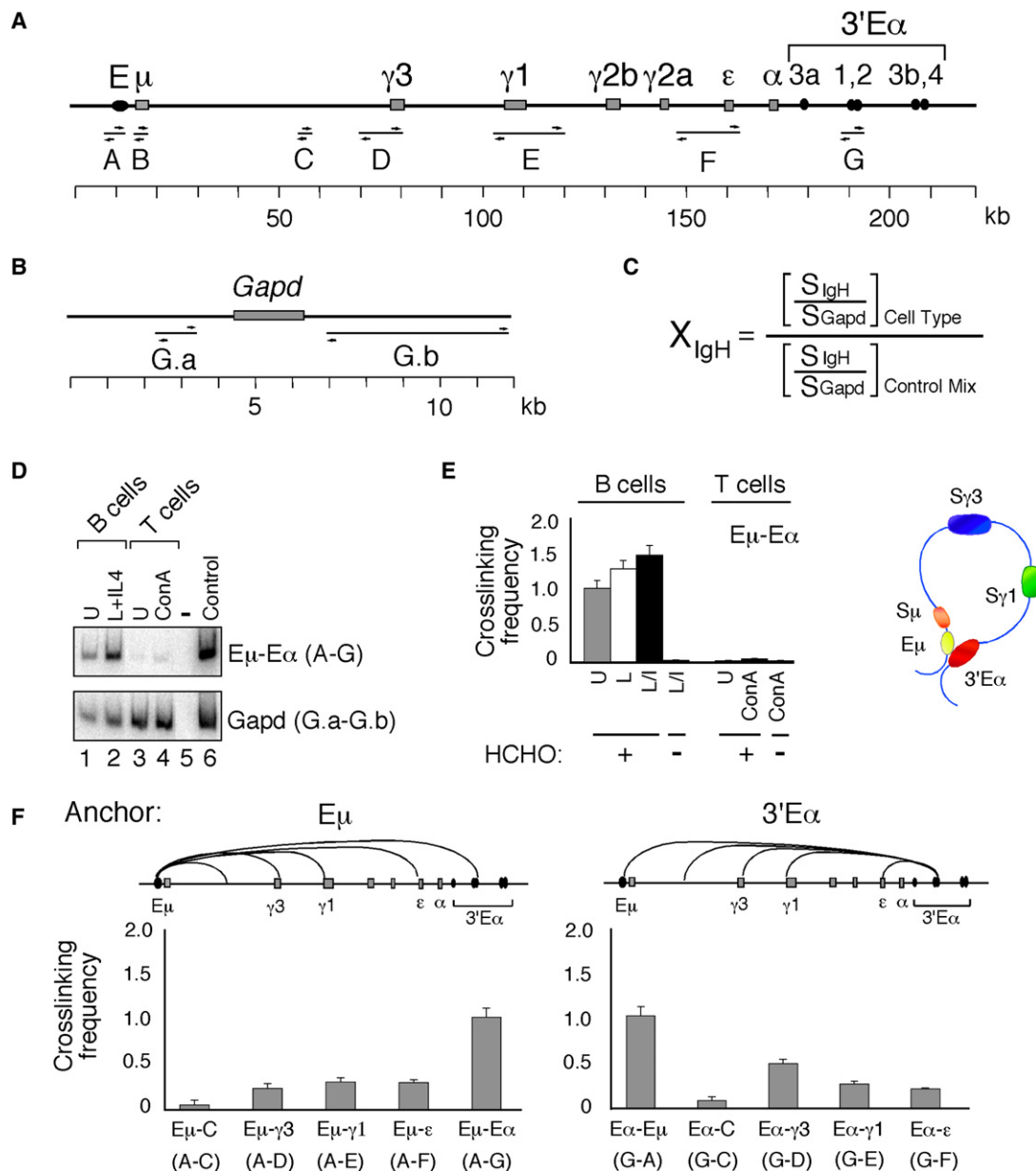


Figure 1. B Cell-Specific Association of the $E\mu$ and $3'E\alpha$ Enhancers and Looping of the Intervening *Igh* Locus

(A) The 3' portion of the *Igh* locus containing C_H genes spans 220 kb (Chevallard et al., 2002). HindIII restriction fragments (A–G) used in the 3C assays are shown. Primers are indicated by arrows above and below each fragment.

(B) The *Gapd* locus is shown, and HindIII restriction fragments G.a and G.b are 3.5 kb apart.

(C) The equation used to calculate the relative crosslinking frequency between two *Igh* restriction fragments (X_{IgH}). S_{IgH} is the signal obtained with primer pairs for two different *Igh* restriction fragments, and S_{Gapd} is the signal obtained with primer pairs for the *Gapd* locus fragments G.a and G.b. This calculation gives the relative crosslinking frequency for each cell type or activation state and corrects for differences in PCR amplification efficiencies, crosslinking and ligation efficiencies, the amounts of template used, and the size of the PCR products.

(D) Representative examples of chromosome conformation capture (3C) assays of the *Igh* locus are shown. Primer pairs for *Igh* locus fragments A and G and *Gapd* fragments G.a and G.b were used in PCR to detect amplification products from crosslinked templates derived from unstimulated (U) and LPS+IL4 (L+I)-stimulated B cells (lanes 1 and 2), unstimulated (U) and ConA-stimulated splenic T cells (lanes 3 and 4), no template (lane 5), and control mix (lane 6). PCR was carried out in the presence of [α - 32 P]dCTP, and products were resolved by 6% PAGE in TBE and then were quantitated by phosphorimaging.

(E) The relative crosslinking frequencies observed in splenic B cells that were unstimulated (U) or activated with LPS (L) or LPS+IL4 (L+IL4) and splenic T cells that were unstimulated (U) or activated with ConA are shown (left). Templates were prepared in the presence or absence of formaldehyde (HCHO) as indicated. A schematic is shown of the looped *Igh* locus in which the $E\mu$ and $3'E\alpha$ enhancers interact (right).

(F) Chromatin templates from unstimulated splenic B cells were analyzed for relative crosslinking frequencies between anchor fragment A ($E\mu$) and fragments C–G (left) and between anchor fragment G ($3'E\alpha$) and fragments A and C–F (right). The average relative crosslinking frequencies were

these interactions are tightly correlated with gene-specific expression (Carter et al., 2002; Tolhuis et al., 2002). Loop structures have also been detected in imprinted (Horike et al., 2005; Murrell et al., 2004) and cytokine (Spilianakis and Flavell, 2004) clusters and between chromatin boundary elements (Blanton et al., 2003). In the *Igh* locus, inducible transcription from the downstream GLT promoters requires the 3'E α LCR (Manis et al., 1998; Pinaud et al., 2001) and suggests that communication might occur through direct interaction between these distant *cis* elements by loop formation. The physical proximity between the downstream GLT promoters with 3'E α may facilitate transcription but would be unlikely to mediate S-S synapsis because the downstream S regions would not be in close proximity to S μ .

By using chromosome conformation capture (3C) techniques, we have examined long-range interactions in the *Igh* locus. In splenic B cells poised for CSR, the *Igh* locus assumed a unique and unanticipated chromosomal loop conformation in vivo in which the E μ enhancer directly interacted with the downstream 3'E α LCR. The S μ region was closely arrayed in *cis* with E μ . The E μ :3'E α structure could facilitate S-S synapsis after GLT promoter activation, because an S region would travel with its proximal GLT promoter to the E μ :3'E α complex and there come into close proximity with S μ . Strong recruitment of GLT promoters to the E μ :3'E α complex was cytokine dependent and correlated with transcription activation. These interactions were dependent on the 3'E α LCR, as indicated by the fact that deletion of the hypersensitive site (hs) 3b,4 largely abrogated E μ :3'E α complex formation and GLT promoter association, whereas deletion of the S μ tandem repeats (TR) had no apparent impact on these LRIs. Deletion of the 220 bp core E μ enhancer had essentially no effect on recruitment of GLT promoters to the enhancer complex and only a slight effect on CSR. Although AID expression was not required for the interaction of E μ with 3'E α , it was required for strong GLT promoter association with the E μ :3'E α complex, indicating that AID stabilizes S-S synapsis either directly through a scaffolding function or indirectly through DNA repair processes emanating from AID-dependent DSB formation in S regions. Our investigation of *Igh* locus-wide intrachromosomal interactions has led to a new model for generating S-S synapsis in which GLT expression is integrally linked to the formation of an architectural scaffold produced through long-range associations between *Igh* transcription regulatory elements.

RESULTS

The Chromosome Conformation Capture Assay for the *Igh* Locus

The chromosome conformation capture (3C) assay (Dekker et al., 2002; Tolhuis et al., 2002) was used to detect

long-range interactions (LRIs) between transcriptional regulatory elements in the C H subregion of the murine *Igh* locus. In this assay, treatment of live cells with formaldehyde leads to crosslinking of proteins to other nearby proteins and DNA. After cleavage with restriction enzyme, intramolecular ligation of interacting crosslinked DNA fragments results when the protein:DNA network is ligated under low concentration conditions. After ligation, crosslinks are reversed and ligation products are quantified by PCR with primer pairs located at the ends of the restriction fragments under study. The crosslinking frequency of any two restriction fragments is proportional to the frequency with which the genomic sites interact and provides a snapshot of the spatial organization of the locus in vivo.

The organization of the C H region of the *Igh* locus is diagrammed in Figure 1A. The E μ intronic enhancer is located at the 5' end of the C H subregion. The 3' regulatory region, 3'E α , contains hypersensitive sites (hs) 3a, hs1,2, hs3b, and hs4 and has been shown to function as an LCR (reviewed in Khamlichi et al., 2000). Further downstream there are additional hs sites 5–7 (Garrett et al., 2005). Sequestered between the two enhancers are eight C H region genes with their attendant S regions and GLT promoters (with the exception of C δ). HindIII restriction fragments were used for 3C analysis of the *Igh* locus (Figure 1A). Fragments A and G contain the E μ and 3'E α enhancers, respectively, whereas fragments B, D, E, and F each encompass an S region with its I exon and GLT promoter. A HindIII fragment with no S region or GLT promoter was used as a negative control (fragment C).

A series of controls to assess the efficacy of the 3C method for the *Igh* locus were performed (see Figures S1–S3 in the Supplemental Data available online). To permit comparison of crosslinking and ligation efficiencies between sample preparations, we measured the crosslinking frequency between two *Gapd* gene-specific HindIII fragments (G.a and G.b), which are separated by 3.5 kb and are located on chromosome 2 (Figure 1B). The equation for calculating the relative crosslinking frequency is shown (Figure 1C). The crosslinking frequency for the two *Gapd* fragments was arbitrarily set to a value of 1 to enable sample comparisons.

The *Igh* Locus Adopts a Looped Configuration in Splenic B Cells but Not in T Cells

We performed 3C analysis on IgM-expressing splenic B and nonexpressing splenic T cells to assess the spatial configuration of the *Igh* locus. In unstimulated B cells, there was an elevated crosslinking frequency for E μ and 3'E α , which are located on separate HindIII fragments (A and G in Figure 1A); such crosslinking was not evident in the absence of chromatin template (Figure 1D, compare lanes 1 and 5). Intrachromosomal association of E μ and 3'E α was cell type specific because little or no

derived from at least three independent chromatin preparations, and each sample was PCR amplified in duplicate at least twice and SEMs are shown. The crosslinking frequency of the two neighboring *Gapd* fragments (3.5 kb apart) is arbitrarily assigned the value of 1. Scaling on the y axis permits direct comparison with all other figures.

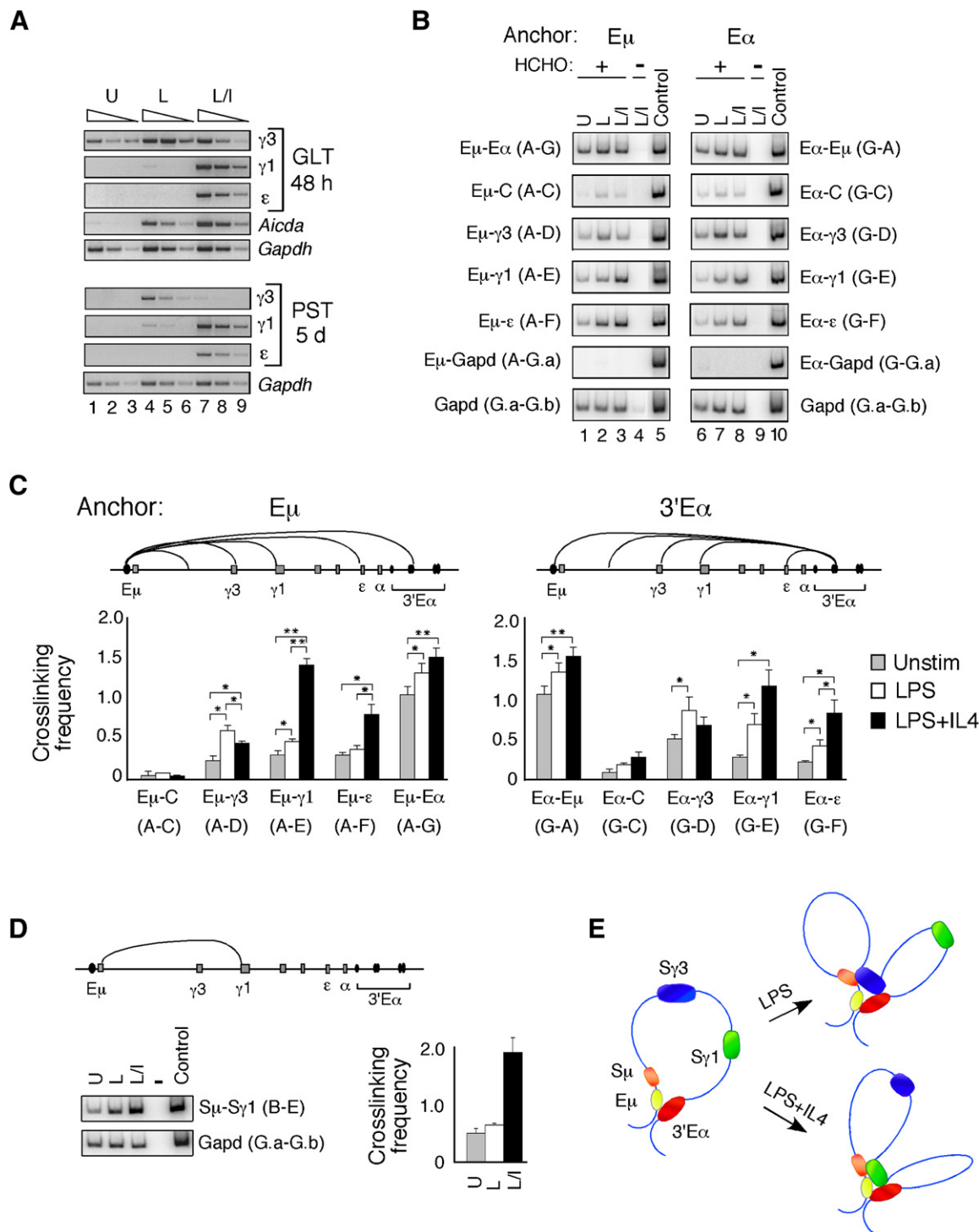


Figure 2. Locus-Wide 3C Assays Indicate that Recruitment of a GLT Promoter to the $E\mu$ -3'E α Enhancers Is Correlated with Transcription Activation

(A) GLTs (top) and post-switch transcripts (PSTs) (bottom) were analyzed by semiquantitative RT-PCR with cDNAs derived from splenic B cells that were unstimulated or activated with LPS or LPS+IL4 for 48 hr (GLTs) and 5 days (PSTs). *Gapdh* PCR products were harvested after 30 cycles (lanes 1, 4, 7), 28 cycles (lanes 2, 5, 8), and 26 cycles (lanes 3, 6, 9). GLT, *Aicda*, and PST PCR products were harvested after 33 cycles (lanes 1, 4, 7), 31 cycles (lanes 2, 5, 8), or 29 cycles (lanes 3, 6, 9).

(B) B cells were unstimulated (U) or activated with LPS (L) or LPS+IL4 (L+I) for 48 hr, and chromatin templates were prepared in the presence or absence of formaldehyde (HCHO) as indicated. Representative examples of crosslinking in B cells between anchors $E\mu$ and 3'E α (fragments A and G) with each other and with all other fragments (C-F) in the *Igh* locus are shown. Crosslinking between each of the anchors $E\mu$ and 3'E α (fragments A and G) with the *Gapd* fragment G.a is shown. Amplification of the control mix containing all combinations of ligated fragments is shown for each primer pair.

amplification was observed in T cells (Figure 1D, lane 3). The absence of E_{μ} and $3'E_{\alpha}$ interaction in T cells does not reflect impaired annealing of primers resulting from polymorphisms because B and T cells were analyzed from the same mouse strain. Consistent results were obtained for three independent chromatin preparations assayed in duplicate in two independent PCR amplifications (Figure 1E). Next, we asked whether the intrachromosomal interaction between E_{μ} and $3'E_{\alpha}$ was altered by cell activation. B cell stimulation with LPS or LPS+IL4 slightly augmented the association of E_{μ} and $3'E_{\alpha}$ and depended on formaldehyde crosslinking, whereas this interaction was not evident in T cells after ConA stimulation (Figures 1D and 1E). These data indicated that the intrachromosomal association of the E_{μ} and $3'E_{\alpha}$ elements created a looped structure encompassing the intervening *Igh* locus in unstimulated and activated splenic B cells (Figure 1E, right). In the nonexpressing T cells, the *Igh* locus was in a linear configuration as evidenced by the absence of E_{μ} and $3'E_{\alpha}$ crosslinking. The association of the E_{μ} and $3'E_{\alpha}$ enhancers in B cells was remarkable, considering that the two sites are separated by 200 kb on the linear chromatin template.

The *Igh* Locus Is Poised for Transcription Activation in B Cells

To further investigate the spatial organization of the *Igh* locus, we undertook a locus-wide assessment of LRIs involving the E_{μ} and $3'E_{\alpha}$ enhancers with other elements of the locus in unstimulated B cells. We found detectable crosslinking between the E_{μ} and $3'E_{\alpha}$ anchor fragments (A and G) and the $\gamma 3$, $\gamma 1$, and ϵ fragments (D, E, and F) that each contain S region and GLT promoter elements (Figure 1F). These intramolecular interactions did not occur due to simple proximity, as shown by the fact that fragment C, which was devoid of S DNA and GLT promoters, was not involved in crosslinking to either the E_{μ} or $3'E_{\alpha}$ enhancer fragments. In addition, crosslinking frequencies were zero for templates isolated from unstimulated splenic T cells (data not shown). These data indicated that the loop created by the intrachromosomal association of E_{μ} and $3'E_{\alpha}$ brought the GLT promoters into sufficient physical proximity with the enhancer core to enable a low amount of crosslinking (Figure 1F). Consequently, in B cells, the enhancers and GLT promoters are in a spatial configuration that is poised for transcription activation and GLT expression.

Long-Range Recruitment of the GLT Promoters to the E_{μ} - $3'E_{\alpha}$ Enhancer Complex Is Transcription Dependent

To relate the spatial configuration of the *Igh* locus and transcriptional status, 3C analyses were performed on splenic B cells that were stimulated to undergo CSR. At 48 hr of B cell activation, GLTs and AID are well expressed and AID-dependent DSBs have formed (Catalan et al., 2003; Rush et al., 2004; Schrader et al., 2005; Wang et al., 2006; Wuerffel et al., 1997), but recombination has not yet transpired (Figure S4; Casellas et al., 1998). The 48 hr time point was used for all 3C analyses because it is likely to be representative of events that are predisposing for CSR whereas later time points may reflect changes that occur as a consequence of recombination. B cell activation by LPS or LPS+IL4 was confirmed by the induction of *Aicda* (AID) and GLT transcription as compared to the *Gapd* loading control (Figure 2A, top). The $\gamma 3$ GLTs were constitutively expressed in unstimulated B cells, induced after LPS activation, and repressed somewhat by the addition of LPS+IL4. In contrast, $\gamma 1$ and ϵ GLT expression were found only after LPS+IL4 induction. CSR was assessed by the post-switch transcript (PST) RT-PCR assay, and switching was detectable at 5 days after activation and correlated with GLT expression (Figure 2B, bottom). The PST switching profile was confirmed by FACS analysis (Figure S5).

Locus-wide 3C studies were performed on LPS- and LPS+IL4-activated B cells (Figure 2B; Figure S6). After LPS activation, the relative crosslinking frequency of E_{μ} enhancer with fragment D, containing the $\gamma 3$ GLT promoter, was 2.5-fold greater than in unstimulated B cells, whereas LPS had little effect on the association of E_{μ} with fragments containing the $\gamma 1$ and ϵ GLT promoters, and these differences were statistically significant (Figure 2C). Conversely, LPS+IL4 treatment slightly reduced the interaction of E_{μ} with the $\gamma 3$ GLT promoter, whereas LRIs between E_{μ} with $\gamma 1$ and ϵ fragments (E, F) were enhanced by 4- and 2.5-fold, respectively, as compared to unstimulated B cells (Figure 2C). Comparable interactions were observed with $3'E_{\alpha}$ as the anchor fragment in locus-wide studies (Figures 2B and 2C). The intrachromosomal interactions detected here were dependent on crosslinking because they did not occur in the absence of formaldehyde treatment, indicating that they did not arise as a result of DNA rearrangements (Figure 2B, lanes 4 and 9; Figure S6). Fragment C and *Gapd* (fragment G.a) showed very little crosslinking with either E_{μ} or $3'E_{\alpha}$ in

(C and D) The average relative crosslinking frequencies were derived from at least three independent chromatin preparations and each sample was PCR amplified in duplicate at least twice and SEMs are shown.

(C) Chromatin templates from unstimulated, LPS-, or LPS+IL4-activated splenic B cells were analyzed for relative crosslinking frequencies between anchor fragment A (E_{μ}) and the rest of the locus fragments (C–G) (left) and between anchor fragment G ($3'E_{\alpha}$) and the rest of the locus fragments (A, C–F) (right). Student's *t* test was used to determine the significance of the different levels of crosslinking frequencies in chromatin samples (**p* < 0.05; ***p* < 0.001).

(D) Chromatin templates from unstimulated, LPS-, or LPS+IL4-activated splenic B cells were analyzed for relative crosslinking frequencies between anchor fragment B (S_{μ}) and fragment E ($\gamma 1$).

(E) A schematic illustrating the long-range interactions between the E_{μ} and $3'E_{\alpha}$ elements of the *Igh* locus in mature B cells. After B cell activation with LPS, the $\gamma 3$ GLT promoter is recruited to the E_{μ} : $3'E_{\alpha}$ complex. Alternatively, LPS+IL4 stimulation triggers the interaction of the $\gamma 1$ GLT promoter with the with the E_{μ} : $3'E_{\alpha}$ enhancer hub.

response to either LPS or LPS+IL4, demonstrating the specificity of these LRIs (Figures 2B and 2C). Furthermore, the relative crosslinking frequency between the S_{μ} fragment (B) and the $\gamma 1$ fragment (E) was elevated 4-fold after LPS+IL4 activation as compared to LPS alone or no treatment, confirming that IL4 specifically affected recruitment of the $\gamma 1$ locus to the region of E_{μ} - S_{μ} (Figure 2D). These studies demonstrated that after the induction of specific GLTs, the active gene was recruited to the E_{μ} -3'E α complex (Figure 2E). This created an ordered spatial architecture that was structured by transcriptional elements and had the effect of bringing the downstream S region targeted for recombination into the vicinity of the E_{μ} - S_{μ} region, thereby promoting S-S proximity. However, these studies did not distinguish the relative contribution of promoter-enhancer interactions from those involved in S-S synopsis.

To determine whether the relative crosslinking frequency at specific isotypes was related to CSR incidence, we analyzed $\mu \rightarrow \gamma 3$ and $\mu \rightarrow \gamma 1$ switching after B cell activation with LPS or LPS+IL4 for 5 days, by FACS analysis (Figure S5). The results from three independent experiments were amalgamated and indicated that $\mu \rightarrow \gamma 3$ and $\mu \rightarrow \gamma 1$ switching occurred in $15\% \pm 2\%$ and $40\% \pm 4\%$ of B cells, respectively. Therefore, the incidence of isotype switching paralleled the relative crosslinking frequency between the E_{μ} enhancer and the relevant GLT promoters, indicating that the interactions between the GLT promoters and the E_{μ} :3'E α complex occurred in switching B cells. It should be noted that the $\gamma 3$ GLT was expressed in unstimulated splenic B cells, providing an explanation for the relatively high association between the $\gamma 3$ GLT promoter and the 3'E α LCR prior to B cell activation.

S Region Sequence Is Not a Major Contributor to LRIs in the *Igh* Locus

The close proximity of the E_{μ} with S_{μ} and the downstream S regions with their paired GLT promoters makes it difficult to discern which elements contribute to the LRIs we detect in the *Igh* locus. To directly assess the importance of S DNA in S-S synopsis, we examined the locus-wide interactions of the *Igh* locus in splenic B cells derived from S_{μ} tandem repeat (TR)-deficient (S_{μ} TR Δ/Δ) mice (Luby et al., 2001). In these mice, all the S_{μ} TRs were removed, with the exception of 15 GAGCT pentamers, which are AID hotspot motifs. CSR frequency was reduced 50%–87% of WT, indicating that the S_{μ} TRs make a relatively modest contribution to the CSR efficiency (Luby et al., 2001). Appropriate B cell activation in S_{μ} TR Δ/Δ mice was achieved with LPS or LPS+IL4 treatment as assessed by the expression of GLTs and the presence of PSTs (Figure S7). Relative crosslinking frequencies between the E_{μ} and 3'E α anchor fragments (A and G) and the downstream $\gamma 3$, $\gamma 1$, and ϵ fragments (D, E, and F) were overall very similar to those found for WT for all stimulation states, and differences in crosslinking frequencies induced by specific activation conditions were statistically significant (p values ranged from $p < 0.05$ to $p < 0.001$) (Figure 3). Fragment C, devoid of a GLT promoter, displayed almost no detectable

interactions with either E_{μ} or 3'E α (Figure 3). Overall, the absence of the S_{μ} TRs has a modest impact on LRIs detected in the *Igh* locus, indicating that S DNA per se was not a major structural element in these associations.

Deletion of the Core E_{μ} Enhancer Modestly Affects GLT Expression, E_{μ} :3'E α Interactions, and CSR

The core E_{μ} (cE_{μ}) enhancer is a 220 bp element that is flanked by two matrix attachment site sequences and is located between the J_H cluster and S_{μ} TRs (reviewed in Ernst and Smale, 1995). VDJ joining is severely reduced in cE_{μ} -deficient (Δ/Δ) mice, and consequently B cells represent only 10%–15% of cells in the spleen (Perlot et al., 2005; Sakai et al., 1999). In contrast, deletion of the cE_{μ} enhancer causes at most a modest reduction of CSR (Perlot et al., 2005; Sakai et al., 1999). To determine whether the cE_{μ} element influences E_{μ} :3'E α complex formation, we analyzed chromatin samples from WT and cE_{μ} Δ/Δ mice, derived from the 129Sve strain, by 3C technology. Resting ($CD43^{-}$) B cells were isolated and were unstimulated or activated with LPS or LPS+IL4, as described above. Real-time RT-PCR analysis of the GLT profile revealed that in cE_{μ} Δ/Δ B cells, the I_{μ} GLT was reduced relative to the WT (Figure S8A), as previously observed for pro-B cells (Perlot et al., 2005). In all other respects, the GLT expression profile appeared nearly identical in WT and cE_{μ} Δ/Δ B cells (Figure S8A). FACS analysis revealed no reduction in $\mu \rightarrow \gamma 1$ switching compared to WT, and $\mu \rightarrow \gamma 3$ switching was reduced 50% of WT in cE_{μ} Δ/Δ B cells (Figure S8B), as expected.

In unstimulated cE_{μ} Δ/Δ B cells, the relative crosslinking frequencies between the E_{μ} and 3'E α fragments (A and G) was reduced to 56% of WT, and this difference is statistically significant ($p = 0.03$), suggesting that cE_{μ} may contribute somewhat to the E_{μ} :3'E α interaction along with other as-yet-undefined elements (Figure 4). This pattern of intrachromosomal interaction was consistent with the reduced I_{μ} GLT expression found in unstimulated cE_{μ} Δ/Δ B cells. After B cell activation, the relative crosslinking frequencies between the E_{μ} and 3'E α fragments (A and G) and the downstream $\gamma 3$, $\gamma 1$, and ϵ fragments (D, E, and F) were overall very similar for both WT and cE_{μ} Δ/Δ samples for all stimulation states (Figure 4). Fragment C, devoid of a GLT promoter, showed almost no interactions with either E_{μ} or 3'E α (Figure 4). We concluded that deletion of the cE_{μ} element modestly influenced the E_{μ} :3'E α intrachromosomal interaction in unstimulated B cells and, upon B cell activation, had little influence on GLT promoter association with the E_{μ} :3'E α complex. The equivalent recruitment of the downstream GLT promoters to the E_{μ} :3'E α complex in WT and cE_{μ} Δ/Δ B cells was strongly correlated with normal amounts of GLTs and CSR.

3'E α Integrity Is Required for Intrachromosomal Interactions in the *Igh* Locus

The 3'E α LCR is responsible for transcription of downstream GLTs, as indicated by the fact that targeted knock-out of *hs3b,4* leads to diminution of GLT expression (with

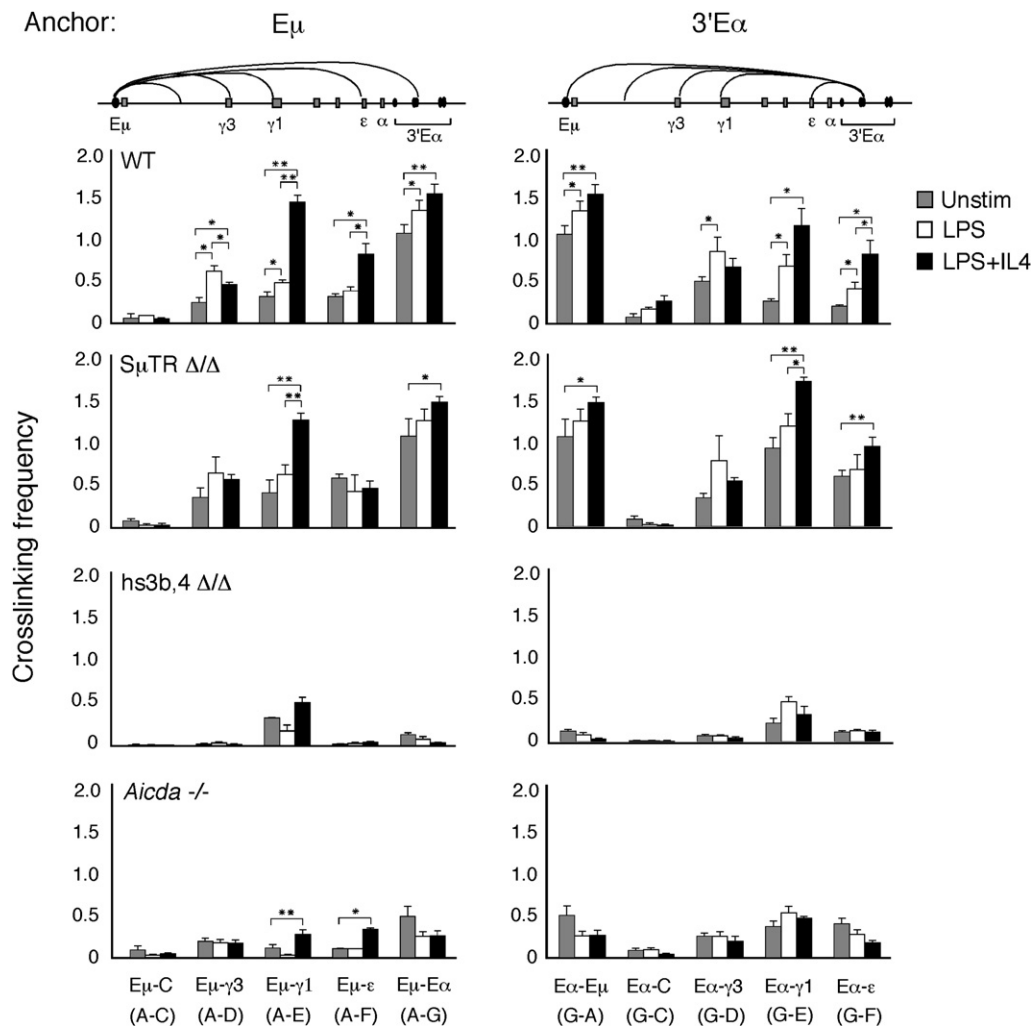


Figure 3. Interaction of $E\mu$ and $3'E\alpha$ with Each Other and with GLT Promoters Does Not Require the $S\mu$ Region but Is Dependent on the Integrity of the $3'E\alpha$ Enhancer and AID Expression

Relative crosslinking frequencies for three to six independent chromatin template preparations are shown for splenic B cells from WT, $S\mu TR \Delta/\Delta$, $3'E\alpha$ $hs3b,4 \Delta/\Delta$, and $Aicda^{-/-}$ mice that were unstimulated or activated with LPS alone or LPS+IL4 for 48 hr. $E\mu$ or $3'E\alpha$ $hs1,2$ were used as anchor fragments that were used together with fragments from the rest of the locus, as indicated. Preparations of templates from C57BL/6 and C57BL/6x129 WT mice gave the same relative crosslinking values, and analyses were amalgamated here. Data represent standard errors of the mean values. Student's t test was used to determine the significance of the different crosslinking frequencies in chromatin samples (*p < 0.05; **p < 0.001).

the exception of $\gamma 1$) (Pinaud et al., 2001). We sought to evaluate the influence of $hs3b,4$ on the ability of the $3'E\alpha$ element to associate with $E\mu$ and GLT promoters by analyzing chromatin templates from the $hs3b,4$ -deficient (Δ/Δ) mouse. The $3'E\alpha$ $hs1,2$ is located on fragment G, whereas $hs3b,4$ is located on a HindIII fragment 7 kb downstream. It was difficult to make primers for many of the HindIII fragments within $3'E\alpha$ because of the reiterated nature of the DNA sequence. Instead, we relied on the proximity of fragment G to $hs3b,4$ as an indicator of the disposition of this element. Analysis of $hs3b,4 \Delta/\Delta$ B cells after activation with LPS and LPS+IL4 showed severely reduced $\gamma 3$ and ϵ GLT and PST expression, whereas $\gamma 1$ GLT and PST production were diminished but still detectable (Figure S7), as previously reported (Pinaud et al.,

2001). In unstimulated $hs3b,4 \Delta/\Delta$ -derived chromatin templates, the relative crosslinking frequency between $E\mu$ and $3'E\alpha$ was reduced to 12% of WT and was not enhanced by LPS or LPS+IL4 treatment (Figure 3).

In stimulated B cells, relative crosslinking frequencies between $E\mu$ and $3'E\alpha$ anchor fragments (A and G) and the downstream $\gamma 3$ and ϵ fragments (D and F) were significantly reduced (p < 0.001) compared to those found for WT for all activation states (Figure 3), demonstrating that the structural integrity of the $E\mu$: $3'E\alpha$ enhancer unit was dependent on the presence of $hs3b,4$ enhancer element and that the $E\mu$: $3'E\alpha$ complex must form to enable subsequent GLT promoter recruitment and transcription from the $\gamma 3$ and ϵ loci. We also observed that although the association of $E\mu$ (fragment A) with the $\gamma 1$ fragment (E) was

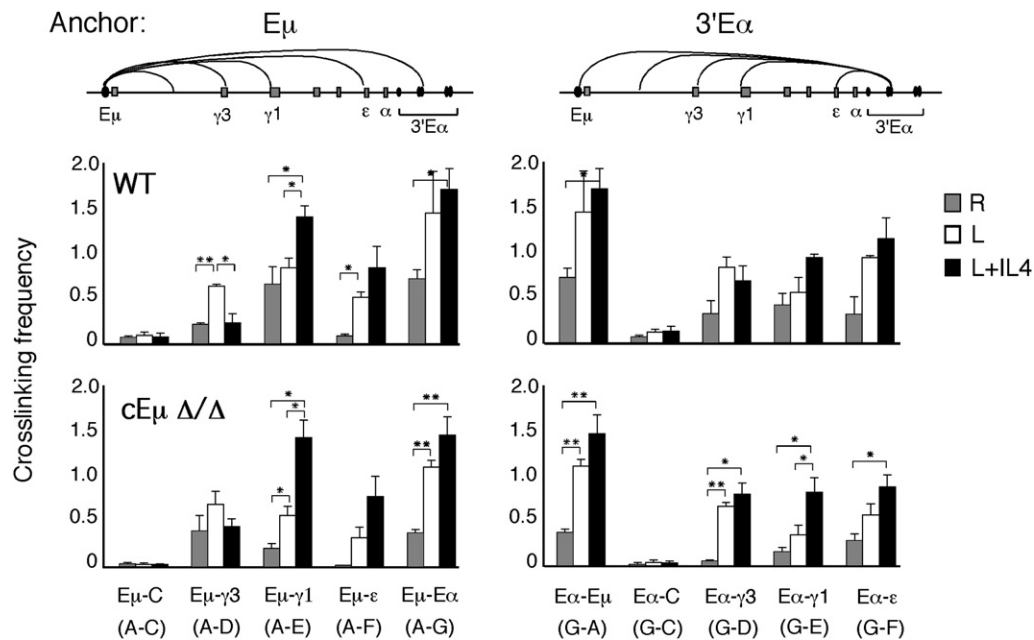


Figure 4. $E\mu:3'E\alpha$ Complex Formation and Recruitment of GLT Promoters Is Not Dependent on the $cE\mu$ Enhancer Element

Relative crosslinking frequencies for two independent chromatin template preparations are shown for $CD43^{-}$ resting splenic B cells (R) from WT and $cE\mu \Delta/\Delta$ mice (129Sve strain) that were unstimulated or activated with LPS alone or LPS+IL4 for 48 hr. $E\mu$ and $3'E\alpha$ $hs1,2$ were used as anchor fragments (A and G) that were used along with fragments from the rest of the locus, as indicated. Data represent standard errors of the mean values. Student's *t* test was used to determine the significance of the different crosslinking frequencies in chromatin samples (* $p < 0.05$; ** $p < 0.001$).

poorly enhanced after B cell activation, the specific IL-4 inducibility of this interaction was retained whereas it was lost for $3'E\alpha$ and $\gamma 1$ fragment interaction. Together, these results demonstrated that in any given $hs3b,4 \Delta/\Delta$ B cell, the $\gamma 1$ locus interacted with either $E\mu$ or $3'E\alpha$ although these elements did not associate with each other. The long-range interaction of $E\mu$ with the $\gamma 1$ fragment preserved the possibility of CSR because $S\mu$ and $S\gamma 1$ were in close proximity, and this was correlated with diminished but detectable $\mu \rightarrow \gamma 1$ switching in $hs3b,4$ -deficient B cells (see Figure S7 and Pinaud et al., 2001).

AID Expression Is a Major Contributor to LRIs in the *Igh* Locus

AID triggers the formation of DSBs in S regions (Rush et al., 2004; Schrader et al., 2005), and these breaks become a focus for recruitment of DNA repair proteins (Celeste et al., 2002; Petersen et al., 2001). To test the proposition that AID contributes to S-S synapsis, we analyzed *Igh* locus-wide interactions in AID-deficient mice. GLT and PST assays, for the *Aicda*^{-/-} B cells, demonstrated appropriate GLT activation but absence of PSTs in response to LPS and LPS+IL4 activation (Figure S7), as previously described (Muramatsu et al., 2000). Surprisingly, in unstimulated *Aicda*^{-/-} B cells, the relative crosslinking frequency between the $E\mu$ and $3'E\alpha$ fragments (A and G) was reduced to 50% of WT (Figure 3). After B cell activation, locus-wide interactions between the $E\mu$ and $3'E\alpha$ anchor fragments (A and G) and the downstream $\gamma 3$, $\gamma 1$, and ϵ fragments (D, E, and F) were reduced to between 14%

and 17% of WT (Figure 3). Nonetheless, intrachromosomal interactions found for *Aicda*^{-/-} B cells are higher than those of $hs3b,4 \Delta/\Delta$ B cells, and these differences were statistically significant ($p < 0.05$) (Figure S9). Strikingly, in AID-deficient B cells, the reduced interaction between the $E\mu$ and $3'E\alpha$ elements and GLT promoters was sufficient for GLT expression. The amount of GLT expression was essentially the same in WT and AID-deficient B cells, so it follows that the additional degree of crosslinking observed in WT chromatin templates arose from AID action independent of transcription. These findings demonstrated a role for AID or processes associated with DSB formation and DNA repair in stabilizing intrachromosomal interactions of the *Igh* locus.

The reduced degree of $E\mu:3'E\alpha$ interaction in AID-deficient unstimulated splenic B cells was unexpected because AID is only slightly expressed in WT cells prior to B cell activation. The low *Aicda* expression in unstimulated cells was highlighted by comparison to that found in LPS+IL4-activated B cells (Figure 5A, compare lanes 1–3 and 7–9). Varying numbers of preactivated B cells were resident in the spleen and expressed *Aicda* and GLTs, whereas these molecules were absent in resting ($CD43^{-}$) WT B cells (Figure 5A). To examine this issue, we sorted for resting ($CD43^{-}$) WT B cells and found that the relative crosslinking frequency between $E\mu$ and $3'E\alpha$ was reduced to 50% of total splenic B cells (Figure 5B). Next, we considered the possibility that the AID expressed in preactivated splenic B cells was responsible for enhanced $E\mu:3'E\alpha$ interaction. The relative crosslinking

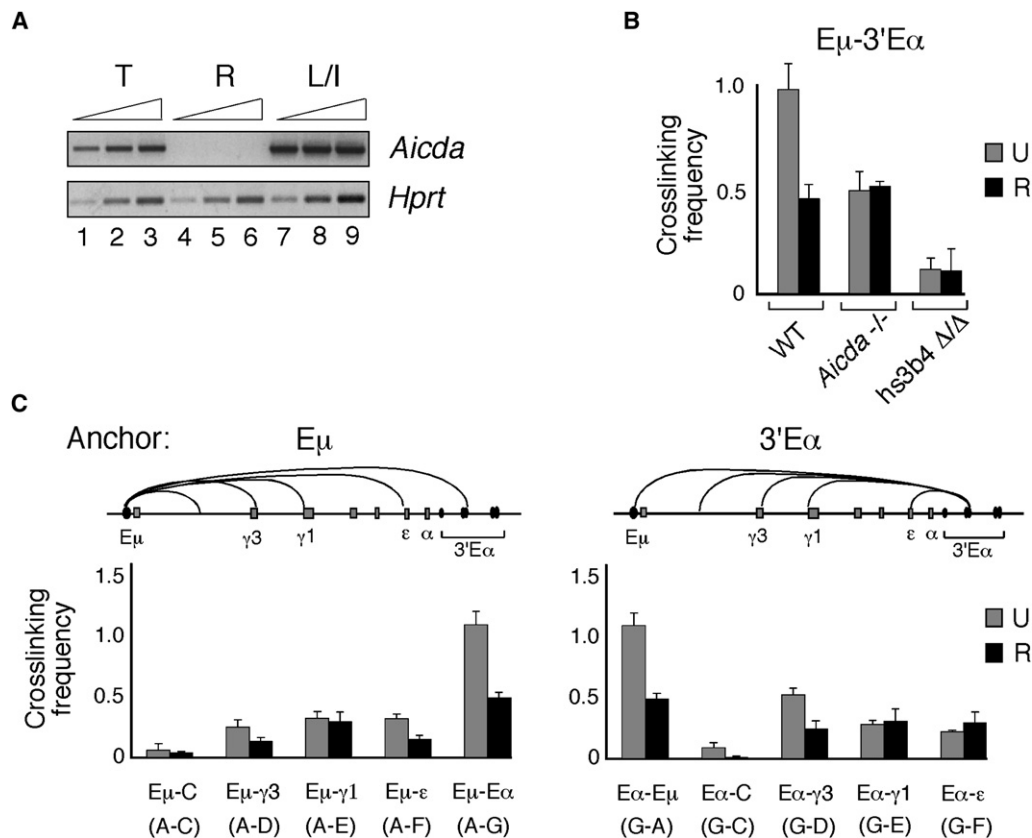


Figure 5. AID Expression Augments the Association of the $E\mu$ Enhancer and 3'E α LCR Elements

Unstimulated splenic B cell preparations (U) were obtained from B cell enrichment columns whereas resting splenic B cells (R) were isolated by sorting for CD43⁺ B cells (Experimental Procedures).

(A) RT-PCR with cDNAs derived from unstimulated total splenic (T) B cells and CD43⁺ resting (R) B cells. *Hprt* PCR products were harvested after 27 cycles (lanes 1, 4, 7), 29 cycles (lanes 2, 5, 8), and 31 cycles (lanes 3, 6, 9). AID PCR products were harvested after 33 cycles (lanes 1, 4, 7), 35 cycles (lanes 2, 5, 8), and 37 cycles (lanes 3, 6, 9).

(B and C) The average crosslinking frequencies along with SEMs were plotted on a scale where a value of 1 corresponds to the crosslinking frequency between the two neighboring *Gapd* fragments, G.a and G.b.

(B) Three independent chromatin template preparations were isolated from WT, *Aicda*^{-/-}, and *hs3b,4* Δ/Δ mice. Chromatin templates were prepared from unstimulated (U) and resting (R) splenic B cells, and crosslinking frequencies were determined for fragments A ($E\mu$) and G (3'E α) and *Gapd* fragments G.a and G.b.

(C) Chromatin templates from unstimulated (U) and CD43⁺ resting (R) splenic B cells from WT mice were analyzed for crosslinking frequencies between anchor fragment A ($E\mu$) and the rest of the *Igh* locus fragments (C-G) (left) and between anchor fragment G (3'E α) and the rest of the locus fragments (A, C-F) (right).

frequencies for $E\mu$ and 3'E α were essentially the same in chromatin templates from *Aicda*^{-/-} resting and total splenic B cells and very similar to WT resting B cells, thereby implicating AID rather than other aspects of B cell activation in stabilizing the $E\mu$:3'E α complex (Figure 5B). We reasoned that AID expression in *hs3b,4* Δ/Δ B cells might enhance $E\mu$:3'E α interaction and obscure the real affect of *hs3b,4* deletion on $E\mu$:3'E α complex formation. Chromatin templates from *hs3b,4* Δ/Δ total and resting B cells showed an essentially identical relative crosslinking frequency for $E\mu$:3'E α and was reduced to 20% of WT and *Aicda*^{-/-} resting B cells (Figure 5B), confirming that $E\mu$:3'E α complex formation was critically dependent on the *hs3b,4* element and leading to the not-mutually-exclusive suggestions that the initial formation of the $E\mu$:3'E α complex was required prior to

AID action and that AID later contributes to the scaffolding integrity.

DISCUSSION

Our studies indicate that the S-S synaptosome is formed in a stepwise process and in a manner contingent upon the presence of *Igh*-specific transcriptional regulatory elements and AID. In mature unstimulated B cells, the distantly located $E\mu$ and 3'E α regulatory elements associate in vivo to form a loop that encompasses the *Igh* C_H genes, and these LRIs are not detected in splenic T cells. After B cell activation, GLT expression is dependent on recruitment of the targeted promoter and flanking S region to the $E\mu$:3'E α complex and, therefore, into close proximity with the S_μ region. The proposition that GLT promoter

and enhancer elements provide a scaffolding for S-S synapsis is supported by the observations that *hs3b,4*, a key regulatory component of the 3'E α , is required for looping of the *Igh* locus as well as recruitment of GLT promoters to the E μ :3'E α enhancer complex. Deletion of the cE μ element leads to a modest reduction in E μ :3'E α interaction and is not required for the recruitment of the GLT promoters to the E μ :3'E α complex in activated B cells, consistent with its modest impact on CSR frequency. Similarly, the S μ TRs are dispensable for these intrachromosomal interactions, and deletion of S μ TRs has a modest effect on CSR. Thus, in B cells poised for CSR, the *Igh* locus assumes a unique and unanticipated chromosomal configuration that creates a structural scaffolding suitable for S-S synaptosome formation.

In unstimulated splenic B cells, looping in the *Igh* locus leads to closer spatial proximity of the GLT promoters with the E μ :3'E α complex to form a "poised" core chromatin conformation that later facilitates GLT expression in response to isotype-specific B cell activation. In the β -globin locus, a poised conformation has been reported in which the β -globin genes are not expressed but promoters and the LCR interact in erythroid cells (Carter et al., 2002; Tolhuis et al., 2002). The chromatin configuration in which a locus is heritably poised for gene expression even when it is transcriptionally silent may be a feature of genes that require rapid transcriptional activation (Smale, 2003). Indeed, B cell stimulation activates rapid GLT expression, consistent with a pre-existing poised *Igh* configuration.

After B cell activation, we found that the long-range interaction of tripartite E μ , 3'E α , and the targeted GLT promoter increased in WT B cells, thus juxtaposing S μ and downstream S regions targeted for transcription. Although these interactions changed only slightly in *Aicda*^{-/-} B cells in response to B cell activation, they were higher than those found in *hs3b,4 Δ / Δ* B cells. Therefore, these intrachromosomal interactions in *Aicda*^{-/-} B cells can account for the presence of GLT expression. However, the relationship between the relative crosslinking levels in 3C assays and gene function is only now being explored, and more work is needed to confirm the significance of the GLT promoter:E μ :3'E α interactions for GLT expression in *Aicda*^{-/-} B cells. Our findings demonstrate that although transcriptional elements provide the architectural basis for initial S-S synapsis, the synaptosome is secondarily stabilized either directly through AID expression or indirectly through recruitment of DNA repair proteins to AID-dependent DSBs in S regions. In support of this interpretation, AID is required to initiate Nbs1 and γ -H2AX repair foci and mutations at S regions during CSR (Petersen et al., 2001). Activated γ -H2AX proteins form foci that spread up to a megabase flanking DSBs and have been implicated in S-S synapsis (Reina-San-Martin et al., 2003; Rogakou et al., 1999). DNA damage-induced foci are centers of DNA repair that simultaneously recruit multiple DSBs (Aten et al., 2004; Lisby et al., 2003), lending weight to the notion that increased S-S synapsis observed in our 3C assays is a consequence of DNA repair proteins coalesced at the sites of AID-induced DNA DSBs. Alternately,

AID may have scaffolding functions associated with its C terminus that mediate the LRLs after B cell activation. AID-directed scaffolding activity is suggested by a naturally occurring deletion at its C terminus that abolishes CSR while retaining the capability for mutagenesis in S μ and V genes and introduction of intra-S μ deletions (Barreto et al., 2003; Ta et al., 2003).

Low-frequency $\mu \rightarrow \gamma$ 1 CSR was recently detected in B cells in which the S μ and S γ 1 regions were replaced by IScel restriction sites, and DSBs were generated by the expression of the IScel enzyme (Zarrin et al., 2006). These observations are consistent with our findings that the *Igh* locus assumes a poised configuration that favors the association of GLT promoters with the E μ :3'E α complex, thereby reducing the distance between S regions, although further work is required to confirm this interpretation. We propose that CSR evolved to ensure that S-S synapsis occurs only as a consequence of AID-dependent DSBs in two targeted S regions. In our model, the S-S synaptosome is formed in a process that couples a unique chromosomal architecture and the introduction of AID-dependent DSBs in two participating S regions directly with DNA repair to reduce the probability of chromosomal translocations.

In sum, our data indicate that the S-S synaptosome is formed as a result of a self-organizing transcription system leading to GLT production. The strict linkage of isotype-specific GLT expression and synapsis between S regions targeted for CSR spatially restricts DNA DSBs and may thereby reduce the danger of genome instability and chromosomal translocations.

EXPERIMENTAL PROCEDURES

Mice, Cell Culture, and FACS Analysis

The 3'E α Δ/Δ mice (Pinaud et al., 2001) and matching wild-type (WT) were provided by J. Manis (Harvard University), S μ TR Δ/Δ mice (Luby et al., 2001) and matching WT were a gift from E. Selsing (Tufts University), and the cE μ Δ/Δ and matched 129Sve WT mice (Perlot et al., 2005) were provided by F. Alt (Harvard University). C57BL/6 \times 129 (AID WT) mice were purchased from Jackson Laboratories. *Aicda*^{-/-} mice (Muramatsu et al., 2000) were a gift from T. Honjo (Kyoto University) and then bred under specific pathogen-free conditions in a fully accredited animal facility at the University of Illinois College of Medicine. Splenic B cells were prepared as described (Shanmugam et al., 2000; Wang et al., 2006) or were sorted for CD43⁺ resting B cells by CD43 magnetic microbeads (MACS, Miltenyi) or were enriched for T cells with the mouse T cell enrichment column kit (MTCC-5, R&D) according to the manufacturer's instructions. B cells were cultured and activated as described (Shanmugam et al., 2000; Wang et al., 2006). Splenic T cells were cultured at a density of 5×10^5 to 1×10^6 and stimulated in RPMI supplemented with 10% FCS and ConcanavalinA (ConA) (5 ng/ml) (15324505, MP Biomedical). Cells were collected at 0 hr, 48 hr, and 5 days to obtain RNA and DNA. Cells from 5 day cultures were washed in HBSS plus 2% FCS and were stained with various antibodies conjugated with fluorescein isothiocyanate (B220 – 553087, IgG3 – 553403, IgG1 – 553443, Thy1.2 – 553003, PharMingen; CD43 – 130-091-585, Miltenyi; IgM – 1020-02, Southern Biotech) and phycoerythrin (B220 – 01125A, PharMingen) or biotin labeled (IgG1 – 553441; PharMingen) and phycoerythrin-cy5 labeled (avidin – 554062; PharMingen). Cells were analyzed with a FACSCalibur with Cellquest software (Becton Dickinson). The flow cytometry analyses represented 5,000–10,000 events and were gated for live

lymphoid cells determined by forward and side scatter and by propidium iodide staining.

RNA Preparation and Semiquantitative RT-PCR

RNA was prepared with TRIzol (GIBCO) according to the manufacturer's instructions. Semiquantitative RT-PCR for GLTs, PSTs, GAPDH, and AID transcripts were performed as described (Wang et al., 2006).

Chromosome Conformation Capture

The chromosome conformation capture (3C) assay (Dekker et al., 2002) was modified for mammalian cells (Tolhuis et al., 2002) and was used with additional adaptations. HindIII (Invitrogen) was used to generate restriction fragments in the *Igh* locus. Chromatin preparations were made from stimulated (5×10^6) and unstimulated (1×10^7) lymphocytes. Splenic B and T cells were washed in Hank's Buffered Saline Solution (HBSS). Cells were crosslinked with 2% formaldehyde for 8 min at room temperature and the reaction was quenched with glycine. Next, nuclei were prepared by incubating the cells in cell membrane lysis buffer (CMLB) (10 mM Tris [pH 7.5], 5 mM MgCl₂, 11% sucrose, 1% Triton X-100) containing protease inhibitors (1 mM PMSF, 10 μ g/ml aprotinin, 10 μ g/ml leupeptin, 10 μ g/ml pepstatinA, 10 μ g/ml phenanthroline, and 16 μ g/ml benzamidin) for 10 min on ice. The cells were pelleted (720 \times g for 10 min at 4°C) and treated again with CMLB. Treatment of the nuclei with HindIII and all subsequent steps, including restriction enzyme inactivation, ligation of the digested crosslinked chromatin, proteinase K treatment, reversal of crosslinks, and DNA purification, were performed as described (Tolhuis et al., 2002). After digestion with HindIII, aliquots of each chromatin sample were analyzed by real-time PCR with primers spanning each HindIII site to assess the degree of digestion (Figure S1). The control template was prepared as essentially described (Tolhuis et al., 2002). In brief, genomic DNA was PCR amplified across each HindIII site of interest to obtain a control fragment that was verified by DNA sequence analysis. DNA fragments spanning the HindIII sites were mixed in equimolar amounts, digested with HindIII, ligated, and added to genomic DNA. A complete laboratory protocol for 3C is available upon request.

PCR Analysis of the Ligation Products

The linear range of PCR amplification was determined for the control template and for the crosslinked chromatin templates by serial dilution as described (Tolhuis et al., 2002). PCR was carried out in 25 μ l reactions containing 200 ng chromatin template DNA, 1.75 mM MgCl₂, 400 μ M each dNTP, 0.3 μ l [α -³²P]dCTP (3000 Ci/mmol), 0.4 μ M each primer, and 1.5 U Platinum Taq (Invitrogen) with initial denaturation for 5 min at 95°C followed by 35 cycles of 94°C for 30 s/58°C for 30 s/72°C for 20 s. Radioactively labeled PCR products were resolved by 6% TBE PAGE and quantitated by phosphorimaging with a STORM 860 and ImageQuant software (Molecular Dynamics). Primer sequences are listed in Figure S3. In general, primers specific for the 3' side of each restriction fragment were used for 3C analysis, with the exception of fragments C and E, where the 5' side was used. All data points were generated from a minimum of three different chromatin samples derived from a minimum of three different mice. Relative crosslinking frequencies were calculated by the equation shown in Figure 1C. Data are presented in histograms as the average crosslinking frequency of all PCR reactions along with the standard error of the mean (SEM).

Supplemental Data

Nine figures and Experimental Procedures are available at <http://www.immunity.com/cgi/content/full/27/5/711/DC1/>.

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